

Sysmex Europe GmbH · Bornbarch 1 · 22848 Norderstedt · Germany

To whom it may concern

Sysmex Europe GmbH

Bornbarch 1 22848 Norderstedt, Germany Phone +49 40 527 26-0 Fax +49 40 527 26-100 info@sysmex-europe.com

LETTER OF AUTHORIZATION

Whereas Sysmex Europe GmbH ("Sysmex"), who are established, reputable and authorised representative in Europe, Africa and Middle East (EMEA region), officially announced by Sysmex Corporation, Japan,

as manufacturer for **Sysmex Coagulation Analyser** with Reagents, Accessories, Software and spare parts and as authorised distributor for **Siemens Coagulation Reagents** in the territory of Moldova (together the "**Products**")

do hereby declare that the company

ECHIPAMED Plus SRL

Valea Trandafirilor 24 "B", off. 80

MD-2001 Chisinau, Moldova (the "COMPANY")

is the non-exclusive distributor of the "Products" in the territory of Moldova.

This declaration is valid until 31 March 2023 and may be revoked unilaterally by Sysmex in writing before that date for due cause. Such due cause shall, among others, be the termination or expiration of the distributorship relationship, if any, between Sysmex and the Company.

On behalf of \$ysmex Europe GmbH

Date: 09 February, 2022

Place: 22848 Norderstedt, Germany

Matthias Voelkel

Senior Executive Officer

sysmex

Sysmex Europe GmbH Bornbarch 1 22848 Norderstedt



Company Location Norderstedt
Registered AG Kiel
HRB 4179
VAT-ID DE 118 687 842
WEEE/ElektroG Reg. Nr. DE 159 56 453

Managing Directors Alain Baverel Seido Biwa Alberto Bonadini Kensuke lizuka Iwane Matsui Stefanie Schaal Jan Willem Schipper Matthias Völkel COMMERZBANK AG, Hamburg IBAN DE20 2004 0000 0287 1879 00 SWIFT/BIC Code COBADEFFXXX



Certificate

Standard

ISO 9001:2015

Certificate Registr. No.

09 100 89004

Certificate Holder:

SYSMEX CORPORATION

1-5-1 Wakinohama-Kaigandori, Chuo-ku, Kobe 651-0073 Japan

including the locations according to annex

Scope:

Development, design, production, sales and servicing of in-vitro diagnostic medical devices, laboratory equipment, reagents and laboratory information system, and development, design, production and sales of customized recombinant protein

Proof has been furnished by means of an audit that the requirements of ISO 9001:2015 are met.

Validity:

The certificate is valid from 2022-05-13 until 2024-07-31. First certification 1998

2022-05-13

MUSICO

TÜV Rheinland Cert GmbH Am Grauen Stein · 51105 Köln





Certificate



Quality Management System EN ISO 13485:2016

Registration No.:

SX 1254782-1

Organization:

SYSMEX CORPORATION

1-5-1 Wakinohama-Kaigandori, Chuo-ku, Kobe

Chuo-ku, Kobe 651-0073 Japan

Scope:

Design and development, manufacture, distribution, installation and service of blood analyzer, urine analyzer, related reagents and accessories Product categories: analyzers and reagents for hematological test, blood coagulation test, immune serum test, biochemical test, genetic test, bacteriological test and urine test

TÜVRheinland

In accordance with EN ISO 13485:2016 Medical devices - Quality management systems - Requirements for regulatory purposes (ISO 13485:2016)

The Certification Body of TÜV Rheinland LGA Products GmbH certifies that the organization has established and applies a quality management system for medical devices.

Proof has been furnished that the requirements specified in the abovementioned standard are fulfilled. The quality management system is subject to yearly surveillance.

Report No .:

150258788-301

Effective date:

2022-04-28

Expiry date:

2024-07-31

Issue date:

2022-04-28

DAKKS

Deutsche
Akkreditierungsstelle
D-ZM-14169-01-02

Michiaki Aihara TÜV Rheinland LGA Products GmbH Tillystraße 2 · 90431 Nürnberg · Germany

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Certificate

Standard

ISO 14001:2015

Certificate Registr. No.

09 104 9374

Certificate Holder:

SYSMEX CORPORATION

1-5-1 Wakinohama-kaigandori, Chuo-ku, Kobe

651-0073, Japan

including the locations according to annex

Scope:

Development, Design, Production, Sales and Servicing Support of In-vitro Diagnostic Medical Devices, Laboratory Equipment, Reagents and Information Technology Systems for Laboratories and Sales of Customized Recombinant Proteins

Proof has been furnished by means of an audit that the requirements of ISO 14001:2015 are met.

Validity:

The certificate is valid from 2020-04-07 until 2023-04-06. First certification 2000

2020-02-25

The stelle

TÜV Rheinland Cert GmbH Am Grauen Stein · 51105 Köln

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(DAKKS

Deutsche Akkreditierungsstelle D-ZM-16031-01-00 TÜVRheinland® Precisely Right. 10/201 10 17 E.Ad. 8 TÜV, TUEV and TLIV are registered trademarks. Utilisation and application requires prior and



To whom it may concern



Konformitätserklärung / Declaration of Conformity

Konformitätserklärung für Siemens Healthcare Diagnostics Products GmbH CE-markierte Produkte.

Hiermit erklären wir, dass ein Konformitätsbeurteilungsverfahren für die hier aufgelisteten In-vitro-Diagnostika-Produkte durchgeführt wurde und sie mit den grundlegenden Anforderungen der Richtlinie 98/79/EG des Europäischen Parlaments und des Rates über In-vitro-Diagnostika übereinstimmen und die Anforderungen gemäß Annex III erfüllt werden.

Declaration of Conformity for Siemens Healthcare Diagnostics Products GmbH CE-marked products.

We hereby declare that a conformity assessment has been performed for the in vitro diagnostic devices listed in the attachment and that they conform to all applicable Essential Requirements of Directive 98/79/EC on in vitro Diagnostic Medical Devices and accordance was shown by conformity assessment procedures of Annex III.

IVD-Kategorie / IVD category:			
Sonstige	Others		
Legaler Hersteller / Legal Manufacturer:			
Siemens Healthcare Dia	gnostics Products GmbH		
Adresse (innerhalb Deutschland):	Address (international):		
Siemens Healthcare Diagnostics Products GmbH Emil-von-Behring-Str. 76 35041 Marburg	Siemens Healthcare Diagnostics Products GmbH Emil-von-Behring-Str. 76 35041 Marburg Germany		

Mit freundlichen Grüßen,

Sincerely,

Siemens Healthcare Diagnostics Products GmbH

Christian Hainer

Regulatory Affairs Manager

Simone Biek

Regulatory Affairs Professional

Datum /Date: 2019-09-17

Anhang /Enclosure: Product List



Konformitätserklärung

Declaration of Conformity

Enclosure to Certification, dd. 2019-09-17

Produktliste | Product List

Product Number (REF)	Package Size	Product Name (English)
	38.00 38.00 38.00 38.00 38.00 38.00 38.00 38.00 38.00 38.00 38.00 38.00 38.00 38.00 38.00 38.00 38.00 38.00 3	

Hemostasis

281007		Thromboclotin	
291070		Dade Ci-Trol 1	
291071		Dade Ci-Trol 2	
291072		Dade Ci-Trol 3	
B4212-40, -50, -100		Dade Innovin	
B4218-1, -2		Dade Actin Activated Cephaloplastin Reagent	
B4218-20, -100		Dade Actin FS Activated PTT Reagent	
B4219-1, -2		Dade Actin FSL Activated PTT Reagent	
B4224-50		Dade Ci-Trol Heparin Control, Low	
B4224-60		Dade Ci-Trol Heparin Control, High	
B4233-15SY		Dade Fibrinogen Determination Reagents	
B4233-22		Dade Data-Fi Abnormal Fibrinogen Control	
B4233-25, -27		Dade Thrombin Reagent	
B4234-25		Dade Owren's Veronal Buffer	
B4238-40		Factor VIII Chromogenic Assay	
B4244-10		Dade Ci-Trol Coagulation Control Level 1	
B4244-20		Dade Ci-Trol Coagulation Control Level 2	
OPAB	03	vWF Ag	
OPAP	03	Protein S Ac	
OPAT	03	PT-Multi Calibrator	
OPBC	03	ProC Ac R	
OPBP	03, 07	INNOVANCE D-Dimer	
OPBR	03	INNOVANCE D-Dimer Sample Diluent	
OPDY	03	INNOVANCE D-Dimer Controls	
OPFH	03, 05	INNOVANCE Antithrombin	
OPHL	03	INNOVANCE VWF Ac	
OQAA	33	Imidazole Buffer Solution	
OQAB	45	Kaolin Suspension	
OQGP	17	LA 1 Screening Reagent	
OQGR	13	LA 2 Confirmation Reagent	
OQGS	29, 35	Pathromtin SL	
OQKE	17	ProC Control Plasma	
OQLS	13	ProC Global	
OQVK	11	Fibrinogen Calibrator Kit	
OQWD	11	LA Control High	



Konformitätserklärung

Declaration of Conformity

Enclosure to Certification, dd. 2019-09-17

OQWE	11	LA Control Low		
OQYG	11	Protein C Reagent		
ORHO	37	Calcium Chloride Solution		
ORKE	41	Control Plasma N		
ORKL	17	Standard Human Plasma		
ORSM	19	Coagulation Factor V Deficient Plasma		
OSDF	13	Coagulation Factor XI Deficient Plasma		
OSDG	13	Coagulation Factor XII Deficient Plasma		
OSGR	13	Coagulation Factor II Deficient Plasma		
OTXV	13	Coagulation Factor VII Deficient Plasma		
OTXW	17	Coagulation Factor VIII Deficient Plasma		
отхх	17	Coagulation Factor IX Deficient Plasma		
OTXY	13	Coagulation Factor X Deficient Plasma		
OUBD 23		Von Willebrand Reagent		
OUBD 37		BC von Willebrand Reagent		
OUBU	15	Berichrom a₂-Antiplasmin		
OUCA	17	Berichrom Plasminogen		
OUHP	29, 49	Thromborel S		
OUIA	15	Berichrom C1-Inhibitor		
OUPZ	17	Control Plasma P		
OUVV	15	Berichrom Protein C		
OWHM	13	Test Thrombin Reagent		
OWNA	11	BC Thrombin Reagent		
OWOA	15	Berichrom PAI		
owsu	11	Berichrom F XIII		
OWWR	15, 17	Berichrom Antithrombin III (A)		
OWZG	19, 23	Multifibren U		

- End of Product List -



EC Declaration of Conformity

Application of Directive	es:
- 98/79/EC of 27 Octob	per 1998 on In Vitro Diagnostic Medical Devices
Means of conformity: The following product - Directive 98/79/EC by Annex III of the Direct	ased on the conformity assessment procedures in accordance with
Product identification: Product name:	CA CLEAN I
Classification:	Other device (except Annex II and self-testing devices)
List of Applied Standard - Harmonised Standards documentation.	rds: s used for conformity assessment are listed in the technical
Legal Manufacturer: Name: Address:	SYSMEX CORPORATION 1-5-1 Wakinohama-Kaigandori, Chuo-ku, Kobe 651-0073, Japan
Authorised officer:	Hiroshi Yamane, Executive Vice President Date: 13 March 2018
Authorised representat	ive:
Name:	SYSMEX EUROPE GMBH

This declaration of conformity is issued under the sole responsibility of the manufacturer and is valid until 25.05.2022 or until a revised declaration is Issued due to product modifications.

Bornbarch 1, 22848 Norderstedt, Germany

Fernando Andreu, Chief Operations Officer



Date: MARCH 21 TOOP

Address:

Authorised officer:



EC Declaration of Conformity

Application of Directives: - 98/79/EC of 27 October 1998 on In Vitro Diagnostic Medical Devices Means of conformity: The following product is in conformity with - Directive 98/79/EC based on the conformity assessment procedures in accordance with Annex III of the Directive. Product identification: Product name: CA CLEAN II Classification: Other device (except Annex II and self-testing devices) List of Applied Standards: - Harmonised Standards used for conformity assessment are listed in the technical documentation. Legal Manufacturer: Name: SYSMEX CORPORATION Address: 1-5-1 Wakinohama-Kaigandori, Chuo-ku, Kobe 651-0073, Japan Authorised officer: Hiroshi Yamane, Executive Vice President Authorised representative: Name: SYSMEX EUROPE GMBH Address: Bornbarch 1, 22848 Norderstedt, Germany

This declaration of conformity is issued under the sole responsibility of the manufacturer and is valid until 25.05.2022 or until a revised declaration is Issued due to product modifications.

Fernando Andreu, Chief Operations Officer

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Date: YEARCH 21 TOIR

Authorised officer:

EC Declaration of Conformity

Application	of	Council	Directive:
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- 98/79/EC of 27 October 1998 on In Vitro Diagnostic Medical Devices

Means of conformity:

The following products are in conformity with

- Directive 98/79/EC based on the conformity assessment procedures in accordance with Annex III,

Product identification:

Product name:

CUVETTE

Model name:

SUC-400A

Classification:

Other device (except Annex II and self-testing devices)

List of Applied Standards:

- Harmonised Standards used for conformity assessment are listed in the technical documentation.

Legal Manufacturer:

Name:

SYSMEX CORPORATION

Address:

1-5-1 Wakinohama-Kaigandori, Chuo-ku, Kobe 651-0073 Japan

Authorised officer:

Hiroshi Yamane Executive Vice President

Authorised representative:

Name:

SYSMEX EUROPE GMBH

Address:

Bornbarch 1, 22848 Norderstedt, Germany

Authorised officer:

Date: MARCH 21 2018

Fernando Andréu, Chief Operations Officer

This declaration of conformity is issued under the sole responsibility of the manufacturer and is valid until 25.05.2022 or until a revised declaration is Issued due to product modifications.

www.s.smex.eg.jp.net